

OVER-COATING AGENT FOR FORMING FINE PATTERNS AND
A METHOD OF FORMING FINE PATTERNS USING SUCH AGENT

BACKGROUND OF THE INVENTION

5 1. Field of the Invention

 This invention relates to an over-coating agent for forming fine patterns in the field of photolithographic technology and a method of forming fine patterns using such agent. More particularly, the invention relates to an over-coating agent for forming or defining fine patterns, such as hole patterns and trench patterns, that can meet today's requirements for higher packing densities and smaller sizes of semiconductor devices.

2. Description of the Related Art

15 In the manufacture of electronic components such as semiconductor devices and liquid-crystal devices, there is employed the photolithographic technology which, in order to perform a treatment such as etching on the substrate, first forms a film (photoresist layer) over the substrate using a so-called radiation-sensitive photoresist which is sensitive to activating radiations, then performs exposure of the film by selective illumination with an activating radiation, performs development to dissolve away the photoresist layer selectively to form an image pattern (photoresist pattern), and forms a variety of patterns including contact providing patterns such as a hole pattern and a trench pattern using the photoresist pattern as a protective layer (mask pattern).

 With the recent increase in the need for higher packing

densities and smaller sizes of semiconductor devices, increasing efforts are being made to form sufficiently fine-line patterns and submicron-electronic fabrication capable of forming patterns with linewidths of no more than 0.20 μm is currently required. As for the activating light rays necessary in the formation of mask patterns, short-wavelength radiations such as KrF, ArF and F_2 excimer laser beams and electron beams are employed. Further, active R&D efforts are being made to find photoresist materials as mask pattern formers that have physical properties adapted to those short-wavelength radiations.

In addition to those approaches for realizing submicron-electronic fabrication which are based on photoresist materials, active R&D efforts are also being made on the basis of pattern forming method with a view to finding a technology that can provide higher resolutions than those possessed by photoresist materials.

For example, JP-5-166717A discloses a method of forming fine patterns which comprises the steps of defining patterns (=photoresist-uncovered patterns) into a pattern-forming resist on a substrate, then coating over entirely the substrate with a mixing generating resist that is to be mixed with said pattern-forming resist, baking the assembly to form a mixing layer on both sidewalls and the top of the pattern-forming resist, and removing the non-mixing portions of said mixing generating resist such that the feature size of the photoresist-uncovered pattern is reduced by an amount comparable to the dimension of said mixing layer. JP-5-241348 discloses a

pattern forming method comprising the steps of depositing a resin, which becomes insoluble in the presence of an acid, on a substrate having formed thereon a resist pattern containing an acid generator, heat treating the assembly so that the
5 acid is diffused from the resist pattern into said resin insoluble in the presence of an acid to form a given thickness of insolubilized portion of the resist near the interface between the resin and the resist pattern, and developing the resist to remove the resin portion through which no acid has
10 been diffused, thereby ensuring that the feature size of the pattern is reduced by an amount comparable to the dimension of said given thickness.

However, in these methods, it is difficult to control the thickness of layers to be formed on the sidewalls of resist patterns. In addition, the in-plane heat dependency of
15 wafers is as great as ten-odd nanometers per degree Celsius, so it is extremely difficult to keep the in-plane uniformity of wafers by means of the heater employed in current fabrication of semiconductor devices and this leads to the problem
20 of occurrence of significant variations in pattern dimensions.

Another approach known to be capable of reducing pattern dimensions is by fluidizing resist patterns through heat treatment and the like. For example, JP-1-307228A discloses a method comprising the steps of forming a resist pattern on a
25 substrate and applying heat treatment to deform the cross-sectional shape of the resist pattern, thereby defining a fine pattern. In addition, JP-4-364021A discloses a method comprising the steps of forming a resist pattern and heating

it to fluidize the resist pattern, thereby changing the dimensions of its resist pattern to form or define a fine-line pattern.

In these methods, the wafer's in-plane heat dependency is only a few nanometers per degree Celsius and is not very problematic. On the other hand, it is difficult to control the resist deformation and fluidizing on account of heat treatment, so it is not easy to provide a uniform resist pattern in a wafer's plane.

An evolved version of those methods is disclosed in JP-7-45510A and it comprises the steps of forming a resist pattern on a substrate, forming a stopper resin on the substrate to prevent excessive thermal fluidizing of the resist pattern, then applying heat treatment to fluidize the resist so as to change the dimensions of its pattern, and thereafter removing the stopper resin to form or define a fine-line pattern. As the stopper resin, a water-soluble resin, specifically, polyvinyl alcohol is employed singly. However, polyvinyl alcohol alone is not highly soluble in water and cannot be readily removed completely by washing with water, introducing difficulty in forming a pattern of good profile. The pattern formed is not completely satisfactory in terms of stability over time. In addition, polyvinyl alcohol cannot be applied efficiently by coating. Because of these and other problems, the method disclosed in JP-7-45510 has yet to be adopted commercially.

JP 2001-281886A discloses a method comprising the steps of covering a surface of a resist pattern with an acidic film

made of a resist pattern size reducing material containing a water-soluble resin, rendering the surface layer of the resist pattern alkali-soluble, then removing said surface layer and the acidic film with an alkaline solution to reduce the feature size of the resist pattern. JP-2002-184673A discloses a method comprising the steps of forming a resist pattern on a substrate, then forming a film containing a water-soluble film forming component on said resist pattern, heat treating said resist pattern and film, and immersing the assembly in an aqueous solution of tetramethylammonium hydroxide, thereby forming a fine-line resist pattern without involving a dry etching step. However, both methods are simply directed to reducing the size of resist trace patterns themselves and therefore are totally different from the present invention in object.

SUMMARY OF THE INVENTION

The present invention has been accomplished in order to solve the aforementioned problems of the prior art and has as an object providing an over-coating agent that can present satisfactory profiles and meet the characteristics required of today's semiconductor devices.

Another object of the invention is to provide a method of forming fine trace patterns using the over-coating agent.

In order to attain the first object, the present invention provides an over-coating agent for forming fine patterns which is applied to cover a substrate having thereon photore-sist patterns and allowed to shrink under heat so that the

spacing between adjacent photoresist patterns is lessened, with the applied film of the over-coating agent being removed substantially completely to form fine patterns, further characterized by containing a copolymer or a mixture of polyvinyl alcohol with a water-soluble polymer other than polyvinyl alcohol.

In order to attain the second object, the present invention provides a method of forming fine patterns comprising the steps of covering a substrate having thereon photoresist patterns with the above-described over-coating agent for forming fine patterns, then applying heat treatment to shrink the applied over-coating agent under the action of heat so that the spacing between adjacent photoresist patterns is lessened, and subsequently removing the applied film of the over-coating agent substantially completely.

In a preferred embodiment, the heat treatment is performed by heating the assembly at a temperature that does not cause thermal fluidizing of the photoresist patterns on the substrate.

DETAILED DESCRIPTION OF THE INVENTION

The present invention is described below in greater below.

The over-coating agent of the invention for forming fine features of patterns is applied to cover a substrate, having photoresist patterns (mask patterns) thereon, including patterns typified by hole patterns or trench patterns, each of these patterns are defined by spacing between adja-

cent photoresist patterns (mask patterns). Upon heating, the applied film of over-coating agent shrinks to increase the width of each of the photoresist patterns, thereby narrowing or lessening adjacent hole patterns or trench patterns as defined by spacing between the photoresist patterns and, thereafter, the applied film is removed substantially completely to form or define fine patterns.

The phrase "removing the applied film substantially completely" as used herein means that after lessening the spacing between adjacent photoresist patterns by the heat shrinking action of the applied over-coating agent, said film is removed in such a way that no significant thickness of the over-coating agent will remain at the interface with the photoresist patterns. Therefore, the present invention does not include methods in which a certain thickness of the over-coating agent is left intact near the interface with the photoresist pattern so that the feature size of the pattern is reduced by an amount corresponding to the residual thickness of the over-coating agent.

The over-coating agent of the invention for forming fine patterns contains a copolymer or a mixture of polyvinyl alcohol (PVA) with a water-soluble polymer other than polyvinyl alcohol.

The water-soluble polymer other than PVA may be any polymer that can dissolve in water at room temperature and various types may be employed without particular limitation; preferred examples include acrylic polymers, vinyl polymers, cellulosic derivatives, alkylene glycol polymers, urea poly-

mers, melamine polymers, epoxy polymers and amide polymers.

Exemplary acrylic polymers include polymers and copolymers having monomeric components, such as acrylic acid, methyl acrylate, methacrylic acid, methyl methacrylate, N,N-dimethylacrylamide, N,N-dimethylaminopropylmethacrylamide, N,N-dimethylaminopropylacrylamide, N-methylacrylamide, diacetone acrylamide, N,N-dimethylaminoethyl methacrylate, N,N-diethylaminoethyl methacrylate, N,N-dimethylaminoethyl acrylate, acryloylmorpholine, etc.

Exemplary vinyl polymers include polymers and copolymers having monomeric components, such as N-vinylpyrrolidone, vinyl imidazolidinone, vinyl acetate, etc.

Exemplary cellulosic derivatives include hydroxypropylmethyl cellulose phthalate, hydroxypropylmethyl cellulose acetate phthalate, hydroxypropylmethyl cellulose hexahydrophthalate, hydroxypropylmethyl cellulose acetate succinate, hydroxypropylmethyl cellulose, hydroxypropyl cellulose, hydroxyethyl cellulose, cellulose acetate hexahydrophthalate, carboxymethyl cellulose, ethyl cellulose, methylcellulose, etc.

Exemplary alkylene glycol polymers include addition polymers and copolymers of ethylene glycol, propylene glycol, etc.

Exemplary urea polymers include those having methylolurea, dimethylolurea, ethyleneurea, etc. as components.

Exemplary melamine polymers include those having methoxymethylated melamine, methoxymethylated isobutoxymethylated melamine, methoxyethylated melamine, etc. as components.

Among epoxy polymers and amide polymers, those which are water-soluble may also be employed.

It is particularly preferred to employ at least one member the group consisting of alkylene glycol polymers, cellululosic derivatives, vinyl polymers and acrylic polymers. Acrylic polymers are most preferred since they provide ease in pH adjustment. Copolymers comprising acrylic polymers and water-soluble polymers other than acrylic polymers are also preferred since during heat treatment, the efficiency of shrinking the spacing between adjacent photoresist patterns (mask patterns) can be increased while maintaining the shape of the photoresist pattern. The water-soluble polymers can be employed either singly or in combination. Exemplary preferred water-soluble polymers include polyacrylate (PAA), polymethacrylate (PMA), and polyvinylpyrrolidone (PVP), etc.

When water-soluble polymers are used as copolymers, the proportions of the components are not limited to any particular values. However, if stability over time is important, the proportion of the acrylic polymer is preferably adjusted to be larger than those of other building polymers. Other than by using excessive amounts of the acrylic polymer, better stability over time can also be obtained by adding acidic compounds such as p-toluenesulfonic acid and dodecylbenzenesulfonic acid.

By using a copolymer or a mixture of PVA with a water-soluble polymer other than PVA as a component of an over-coating agent, the shrinkage amount (the amount of heat shrinking) of the over-coating agent at a given heating tem-

perature can be increased and thus fine-line patterns can be more efficiently formed in the present invention compared with the conventional cases.

From the viewpoint of increasing the shrinkage amount,
5 it is preferred in the over-coating agent according to the present invention that PVA is copolymerized or mixed in an amount of 0.1 - 5 times by weight, still preferably 0.2 - 2 times by weight, as much as the water-soluble polymer other than PVA.

10 In addition to the copolymer or the mixture of PVA with a water-soluble polymer other than PVA, other polymer(s) or mixtures(s) of water-soluble polymers may be contained. For example, use may be made of a blend of PVA/PAA copolymer with a PAA/PVP copolymer.

15 The over-coating agent for forming fine patterns may additionally contain water-soluble amines. For special purposes such as preventing the generation of impurities and pH adjustment, water-soluble amines that have pKa (acid dissociation constant) values of 7.5 - 13 in aqueous solution at 25 °C
20 are preferably used. Specific examples include the following: alkanolamines, such as monoethanolamine, diethanolamine, triethanolamine, 2-(2-aminoethoxy)ethanol, N,N-dimethylethanolamine, N,N-diethylethanolamine, N,N-dibutylethanolamine, N-methylethanolamine, N-
25 ethylethanolamine, N-butylethanolamine, N-methyldiethanolamine, monoisopropanolamine, diisopropanolamine and triisopropanolamine; polyalkylenepolyamines, such as diethylenetriamine, triethylenetetramine, propylenediamine,

N,N-diethylethylenediamine, 1,4-butanediamine, N-ethyl-
ethylenediamine, 1,2-propanediamine, 1,3-propanediamine and
1,6-hexanediamine; aliphatic amines, such as triethylamine,
2-ethyl-hexylamine, dioctylamine, tributylamine, tripro-
5 pylamine, triallylamine, heptylamine and cyclohexylamine;
aromatic amines, such as benzylamine and diphenylamine; and
cyclic amines, such as piperazine, N-methyl-piperazine and
hydroxyethylpiperazine. Preferred water-soluble amines are
those having boiling points of 140 °C (760 mmHg) and above, as
10 exemplified by monoethanolamine and triethanolamine.

If the water-soluble amine is to be added, it is pref-
erably incorporated in an amount of about 0.1 - 30 mass%,
more preferably about 2 - 15 mass%, of the over-coating agent
(in terms of solids content). If the water-soluble amine is
15 incorporated in an amount of less than 0.1 mass%, the coating
fluid may deteriorate over time. If the water-soluble amine
is incorporated in an amount exceeding 30 mass%, the photore-
sist pattern being formed may deteriorate in shape.

For such purposes as reducing the dimensions of pat-
20 terns and controlling the occurrence of defects, the over-
coating agent for forming fine patterns may further option-
ally contain non-amine based, water-soluble organic solvents.

As such non-amine based, water-soluble organic solvents,
any non-amine based organic solvents that can mix with water
25 may be employed and they may be exemplified by the following:
sulfoxides, such as dimethyl sulfoxide; sulfones, such as di-
methylsulfone, diethylsulfone, bis(2-hydroxyethyl)sulfone and
tetramethylenesulfone; amides, such as N,N-dimethylformamide,

N-methylformamide, N,N-dimethylacetamide, N-methylacetamine and N,N-diethylacetamide; lactams, such as N-methyl-2-pyrrolidone, N-ethyl-2-pyrrolidone, N-propyl-2-pyrrolidone, N-hydroxymethyl-2-pyrrolidone and N-hydroxyethyl-2-

5 pyrrolidone; imidazolidinones, such as 1,3-dimethyl-2-imidazolidinone, 1,3-diethyl-2-imidazolidinone and 1,3-diisopropyl-2-imidazolidinone; and polyhydric alcohols and derivatives thereof, such as ethylene glycol, ethylene glycol monomethyl ether, ethylene glycol monoethyl ether, ethylene
10 glycol monobuthyl ether, ethylene glycol monomethyl ether acetate, ethylene glycol monoethyl ether acetate, diethylene glycol, diethylene glycol monomethyl ether, diethylene glycol monoethyl ether, diethylene glycol monobuthyl ether, propylene glycol, propylene glycol monomethyl ether, glycerol, 1,2-
15 butylene glycol, 1,3-butylene glycol and 2,3-butylene glycol. Among those mentioned above, polyhydric alcohols and their derivatives are preferred for the purposes of reducing the dimensions of patterns and controlling the occurrence of defects and glycerol is particularly preferred. The non-amine
20 based, water-soluble organic solvents may be used either singly or in combination.

If the non-amine based, water-soluble organic solvent is to be added, it is preferably incorporated in an amount of about 0.1 - 30 mass%, more preferably about 0.5 - 15 mass%,
25 of the water-soluble polymer. If the non-amine based, water-soluble organic solvent is incorporated in an amount of less than 0.1 mass%, its defect reducing effect tends to decrease. Beyond 30 mass%, a mixing layer is liable to form at the in-

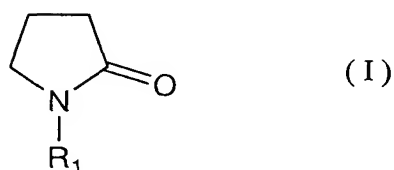
terface with the photoresist pattern.

In addition, the over-coating agent may optionally contain a surfactant for attaining special effects such as coating uniformity and wafer's in-plane uniformity.

5 The surfactant is preferably employed that, when added to the copolymer or the mixture of PVA with a water-soluble polymer other than PVA, exhibits certain characteristics such as high solubility, non-formation of a suspension and miscibility with the polymer component. By using surfactants that
10 satisfy these characteristics, the occurrence of defects can be effectively controlled that is considered to be pertinent to microforming upon coating the over-coating agent.

From the points above, surfactants in the invention are preferably employed at least the one selected among N-
15 alkylpyrrolidones, quaternary ammonium salts and phosphate esters of polyoxyethylene.

N-alkylpyrrolidones as surfactant are preferably represented by the following general formula (I):

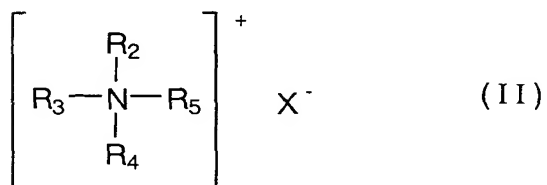


20 where R₁ is an alkyl group having at least 6 carbon atoms.

Specific examples of N-alkylpyrrolidones as surfactant include N-hexyl-2-pyrrolidone, N-heptyl-2-pyrrolidone, N-octyl-2-pyrrolidone, N-nonyl-2-pyrrolidone, N-decyl-2-pyrrolidone, N-undecyl-2-pyrrolidone, N-dodecyl-2-pyrrolidone,
25 N-tridecyl-2-pyrrolidone, N-tetradecyl-2-pyrrolidone, N-

pentadecyl-2-pyrrolidone, N-hexadecyl-2-pyrrolidone, N-heptadecyl-2-pyrrolidone and N-octadecyl-2-pyrrolidone. Among these, N-octyl-2-pyrrolidone ("SURFADONE LP 100" of ISP Inc.) is preferably used.

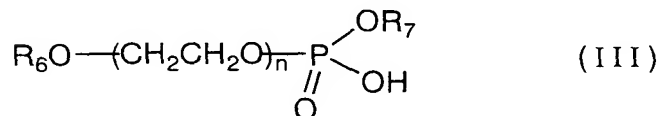
5 Quaternary ammonium salts as surfactant are preferably represented by the following general formula (II):



where R_2 , R_3 , R_4 and R_5 are each independently an alkyl group or a hydroxyalkyl group (provided that at least one of them
10 is an alkyl or hydroxyalkyl group having not less than 6 carbon atoms); X^- is a hydroxide ion or a halogenide ion.

Specific examples of quaternary ammonium salts as surfactant include dodecyltrimethylammonium hydroxide, tridecyltrimethylammonium hydroxide, tetradecyltrimethylammonium hydroxide, pentadecyltrimethylammonium hydroxide, hexadecyl-
15 trimethylammonium hydroxide, heptadecyltrimethylammonium hydroxide and octadecyltrimethylammonium hydroxide. Among these, hexadecyltrimethylammonium hydroxide is preferably used.

Phosphate esters of polyoxyethylene are preferably represented by the following general formula (III):
20



where R_6 is an alkyl or alkylaryl group having 1-10 carbon atoms; R_7 is a hydrogen atom or $(CH_2CH_2O)R_6$ (where R_6 is as de-

lined above); n is an integer of 1 - 20.

To mention specific examples, phosphate esters of polyoxyethylene that can be used as surfactants are commercially available under trade names "PLYSURF A212E" and "PLYSURF

5 A210G" from Dai-ichi Kogyo Seiyaku Co., Ltd.

If the surfactant is to be added, it is preferably incorporated in an amount of about 0.1 - 10 mass%, more preferably about 0.2 - 2 mass%, of the over-coating agent (in terms of solids content). By adopting the amount as described
10 above ranges, it may effectively prevent the variations in the percent shrinkage of patterns, potentially depending on the wafer's in-plane uniformity which is caused by the deterioration of coating property, and also prevent the occurrence of defects that are considered to have cause-and-effect
15 relations with microfoaming on the applied film that generates as the coating conditions are worsened.

The over-coating agent of the invention for forming fine patterns is preferably used as an aqueous solution at a concentration of 3 - 50 mass%, more preferably at 5 - 30
20 mass%. If the concentration of the aqueous solution is less than 3 mass%, poor coverage of the substrate may result. If the concentration of the aqueous solution exceeds 50 mass%, there is no appreciable improvement in the intended effect that justifies the increased concentration and the solution
25 cannot be handled efficiently.

As already mentioned, the over-coating agent of the invention for forming fine patterns is usually employed as an aqueous solution using water as the solvent. A mixed solvent

system comprising water and an alcoholic solvent may also be employed. Exemplary alcoholic solvents are monohydric alcohols including methyl alcohol, ethyl alcohol, propyl alcohol and isopropyl alcohol. These alcoholic solvents are mixed
5 with water in amounts not exceeding about 30 mass%.

The over-coating agent of the invention for forming fine patterns has the advantage of improving resolution beyond the values inherent in photoresist materials and it can attain wafer's in-plane uniformity by eliminating the pattern
10 variations in the plane of the substrate. Further, the over-coating agent of the invention can form patterns of good profile by eliminating the irregularities (roughness) in the shape of patterns due, for example, to the reflection of fluorescent light from the substrate. Yet another advantage
15 of the over-coating agent is its ability to effectively increase the shrinkage amount (the amount of heat shrinking) at a given heating temperature, thereby achieving a remarkably improved effect of forming fine-line patterns.

The method of forming fine-line patterns according to
20 the second aspect of the invention comprises the steps of covering a substrate having photoresist patterns thereon with the above-described over-coating agent for forming fine patterns, then applying heat treatment to shrink the applied over-coating agent under the action of heat so that the spacing
25 between adjacent photoresist patterns is reduced, and subsequently removing the applied film of the over-coating agent substantially completely.

The method of preparing the substrate having photore-

sist patterns thereon is not limited to any particular type and it can be prepared by conventional methods employed in the fabrication of semiconductor devices, liquid-crystal display devices, magnetic heads and microlens arrays. In an exemplary method, a photoresist composition of chemically amplifiable or other type is spin- or otherwise coated on a substrate such as a silicon wafer and dried to form a photoresist layer, which is illuminated with an activating radiation such as ultraviolet, deep-ultraviolet or excimer laser light through a desired mask pattern using a reduction-projection exposure system or subjected to electron beam photolithography, then heated and developed with a developer such as an alkaline aqueous solution, typically a 1-10 mass% tetramethylammonium hydroxide (TMAH) aqueous solution, thereby forming a photoresist pattern on the substrate.

The photoresist composition serving as a material from which photoresist patterns are formed is not limited in any particular way and any common photoresist compositions may be employed including those for exposure to i- or g-lines, those for exposure with an excimer laser (e.g. KrF, ArF or F₂) and those for exposure to EB (electron beams).

[a.] Over-coating agent application step

After thusly forming the photoresist pattern as a mask pattern, the over-coating agent for forming fine patterns is applied to cover entirely the substrate. After applying the over-coating agent, the substrate may optionally be pre-baked at a temperature of 80 - 100 °C for 30 - 90 seconds.

The over-coating agent may be applied by any methods

commonly employed in the conventional heat flow process. Specifically, an aqueous solution of the over-coating agent for forming fine patterns is applied to the substrate by any known application methods including bar coating, roll coating
5 and whirl coating with a spinner.

[b.] Heat treatment (thermal shrinkage) step

In the next step, heat treatment is performed to cause thermal shrinkage of the film of the over-coating agent. Under the resulting force of thermal shrinkage of the film, the
10 dimensions of the photoresist pattern in contact with the film will increase by an amount equivalent to the thermal shrinkage of the film and, as the result, the photoresist pattern widens and accordingly the spacing between adjacent photoresist patterns lessens. The spacing between adjacent
15 photoresist patterns determines the diameter or width of the pattern elements to be finally obtained, so the decrease in the spacing between adjacent photoresist patterns contributes to reducing the diameter of each element of a hole pattern or the width of each element of a trench pattern, eventually
20 leading to the definition of a pattern with smaller feature sizes.

The heating temperature is not limited to any particular value as long as it is high enough to cause thermal shrinkage of the film of the over-coating agent and form or
25 define a fine pattern. Heating is preferably done at a temperature that will not cause thermal fluidizing of the photoresist pattern. The temperature that will not cause thermal fluidizing of the photoresist pattern is such a temperature

that when a substrate on which the photoresist pattern has been formed but no film of the over-coating agent has been formed is heated, the photoresist pattern will not experience any dimensional changes. Performing a heat treatment under
5 such temperature conditions is very effective for various reasons, e.g. a fine-line pattern of good profile can be formed more efficiently and the duty ratio in the plane of a wafer, or the dependency on the spacing between photoresist patterns in the plane of a wafer, can be reduced. Considering
10 the softening points of a variety of photoresist compositions employed in current photolithographic techniques, the preferred heat treatment is usually performed within a temperature range of about 80 - 160 °C for 30 - 90 seconds, provided that the temperature is not high enough to cause thermal flu-
15 idizing of the photoresist.

The thickness of the film of the over-coating agent for the formation of fine-line patterns is preferably just comparable to the height of the photoresist pattern or high enough to cover it.

20 [c.] Over-coating agent removal step

In the subsequent step, the remaining film of the over-coating agent on the patterns is removed by washing with an aqueous solvent, preferably pure water, for 10 - 60 seconds. Prior to washing with water, rinsing may optionally be per-
25 formed with an aqueous solution of alkali (e.g. tetramethylammonium hydroxide (TMAH) or choline). The over-coating agent of the present invention is easy to remove by washing with water and it can be completely removed from the substrate and

the photoresist pattern.

As a result, each pattern on the substrate has a smaller feature size because each pattern is defined by the narrowed spacing between the adjacent widened photoresist patterns.

The fine-line pattern thus formed using the over-coating agent of the present invention has a pattern size smaller than the resolution limit attainable by the conventional methods. In addition, it has a good enough profile and physical properties that can fully satisfy the characteristics required of semiconductor devices.

Steps [a.] - [c.] may be repeated several times. By repeating steps [a.] - [c.] several times, the photoresist trace patterns (mask patterns) can be progressively widened.

The technical field of the present invention is not limited to the semiconductor industry and it can be employed in a wide range of applications including the fabrication of liquid-crystal display devices, the production of magnetic heads and even the manufacture of microlens arrays.

EXAMPLES

The following examples are provided for further illustrating the present invention but are in no way to be taken as limiting. Unless otherwise noted, all amounts of ingredients are expressed in mass%.

EXAMPLE 1

A copolymer of PVA and PAA (2 g; "KURASTMER AP20", product of Kuraray Co., Ltd.), triethanolamine (0.18 g) and a

polyoxyethylenephosphate ester surfactant (0.02 g; "PLYSURF A210G", product of Dai-ichi Kogyo Seiyaku Co, Ltd.) were dissolved in water (52 g) to prepare an over-coating agent.

A substrate was whirl coated with a positive-acting
5 photoresist TDUR-P036PM (product of Tokyo Ohka Kogyo Co., Ltd.) and baked at 80 °C for 90 seconds to form a photoresist layer in a thickness of 0.48 μm.

The photoresist layer was exposed with an exposure unit (Canon FPA-3000EX3, product of Canon Inc.), subjected to heat
10 treatment at 120 °C for 90 seconds and developed with an aqueous solution of 2.38 mass% TMAH (tetramethylammonium hydroxide) to form photoresist patterns which defined hole patterns with an each diameter of 178.9 nm (i.e., the spacing between the photoresist patterns, or the initial hole dimension, was
15 178.9 nm).

The previously prepared over-coating agent was applied onto the substrate including hole patterns and subjected to heat treatment at 115 °C for 60 seconds, thereby reducing the each size of the hole patterns. Subsequently, the substrate
20 was brought into contact with pure water at 23 °C to remove the over-coating agent. The each diameter of the hole patterns was reduced to 152.2 nm.

EXAMPLE 2

A copolymer of PVA and PAA (1 g; "KURASTMER AP20",
25 product of Kuraray Co., Ltd.), a copolymer of PAA and PVP [1 g; PAA:PVP = 2:1 (polymerization ratio)], and a polyoxyethylenephosphate ester surfactant (0.02 g; "PLYSURF A210G", product of Dai-ichi Kogyo Seiyaku Co, Ltd.) were dissolved in

water (33 g) to prepare an over-coating agent.

A substrate was whirl coated with a positive-acting photoresist TDUR-P036PM (product of Tokyo Ohka Kogyo Co., Ltd.) and baked at 115 °C for 90 seconds to form a photoresist layer in a thickness of 0.48 μm.

The photoresist layer was exposed with an exposure unit (Canon FPA-3000EX3, product of Canon Inc.), subjected to heat treatment at 120 °C for 90 seconds and developed with an aqueous solution of 2.38 mass% TMAH (tetramethylammonium hydroxide) to form photoresist patterns which defined hole patterns with an each diameter of 178.9 nm (i.e., the spacing between the photoresist patterns, or the initial hole dimension, was 178.9 nm).

The previously prepared over-coating agent was applied onto the substrate including hole patterns and subjected to heat treatment at 115 °C for 60 seconds, thereby reducing the each size of the hole patterns. Subsequently, the substrate was brought into contact with pure water at 23 °C to remove the over-coating agent. The each diameter of the hole patterns was reduced to 149.8 nm.

COMPARATIVE EXAMPLE 1

A copolymer of PAA and PVP [2 g; PAA:PVP = 2:1 (polymerization ratio)] and a polyoxyethylenephosphate ester surfactant (0.02 g; "PLYSURF A210G", product of Dai-ichi Kogyo Seiyaku Co, Ltd.) were dissolved in water (33 g) to prepare an over-coating agent.

The thusly prepared over-coating agent was then applied onto the substrate including hole patterns that were formed

in the same manner as described in Example 1 (the initial hole dimension, was 178.9 nm), and subjected to heat treatment at 115 °C for 60 seconds, thereby reducing the each size of the hole patterns. Subsequently, the substrate was brought
5 into contact with pure water at 23 °C to remove the over-coating agent. The each diameter of the hole patterns was 158.9 nm.

COMPARATIVE EXAMPLE 2

The same procedure as described in Example 1 was re-
10 peated, except that a homopolymer of PVA (2 g) was used instead of the copolymer of PVA and PAA (2 g; "KURASTMER AP20", product of Kuraray Co., Ltd.). A visual check revealed that residues were remained on the hole patterns due to the insufficient solubility in water of the homopolymer of PVA.

15 As described above in detail, the over-coating agent of the present invention can effectively increase a shrinkage amount (the amount of heat shrinking) in a heating treatment, thereby achieving a remarkably improved effect of forming or defining fine-line patterns.